Rules of **Department of Natural Resources**

Division 60—Public Drinking Water Program Chapter 5—Laboratory and Analytical Requirements

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Title 10—DEPARTMENT OF NATURAL RESOURCES

Division 60—Public Drinking Water Program Chapter 5—Laboratory and Analytical Requirements

10 CSR 60-5.010 Accepted and Alternate Procedures for Analyses

PURPOSE: This rule lists manuals containing acceptable analysis procedures for determination of contaminant levels.

PUBLISHER'S NOTE: The publication of the full text of the material that the adopting agency has incorporated by reference in this rule would be unduly cumbersome or expensive. Therefore, the full text of that material will be made available to any interested person at both the Office of the Secretary of State and the office of the adopting agency, pursuant to section 536.031.4, RSMo. Such material will be provided at the cost established by state law.

(1) Inorganic and Radiological Contaminants. Unless substitute methods are approved by the department, analysis shall be conducted in accordance with the methods listed in the following table. All analytical methods listed in this rule are incorporated by reference. Criteria for analyzing arsenic, barium, beryllium, cadmium, calcium, chromium, copper, lead, nickel, selenium, sodium, and thallium with digestion or directly without digestion, and other analytical test procedures are contained in "Technical Notes on Drinking Water Methods," EPA-600/R-94-173, October 1994. This document also contains approved analytical test methods which remain available for compliance monitoring until July 1, 1996. These methods will not be available for use after July 1, 1996. This document is available from the National Technical Information Service, NTIS PB95-104766, U.S. Department of Commerce, 5285 Port Royal Road, Springfield, VA 22161. The toll-free number is 800-553-6847.

Contaminant	Methodology	Method	Reference
	(if appropriate)	(if appropriate)	
luminum	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
iaiiiiaiii	inductively coupled I lasina—Emission spectroscopy	3120 B	17
	Industrials Counted Disams Mass Constrainment		22
	Inductively Coupled Plasma—Mass Spectrometry	200.8	
	Atomic Absorption—Platform Technique	200.9	22
		3113 B	17
		3111 D	17
ntimony	Atomic Absorption—Gaseous Hydride	D3697-92	19
	Atomic Absorption—Graphite Furnace	3113 B	17
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
	Atomic Absorption—Platform Technique	200.8	22
	•		
rsenic	Atomic Absorption—Graphite Furnace	D2972-93C	19
		3113 B	17
	Atomic Absorption—Gaseous Hydride	D2972-93B	19
	, J	3114 B	17
	Atomic Absorption—Platform	200.9	22
	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
	inductively Coupled Plasma—Emission Spectroscopy		
		3120 B	17
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
sbestos	Transmission Electron Microscopy	100.1	7
	17	100.2	27
arium	Industivaly Counted Diagna, Mass Speatrometry	200.8	22
arrum	Inductively Coupled Plasma—Mass Spectrometry		
	Atomic Absorption—Graphite Furnace	3113 B	17
	Atomic Absorption—Direct Aspiration	3111 D	17
	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
		3120 B	17
eryllium	Atomic Absorption—Graphite Furnace	D3645-93B	19
)		3113 B	17
	Atomic Absorption—Platform	200.9	22
	Inductively Coupled Plasma—Emission Spectroscopy	3120 B	17
		200.7	22
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
admium	Atomic Absorption—Graphite Furnace	3113 B	17
	Atomic Absorption—Platform	200.9	22
	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
	Inductively Coupled Plasma—Emission spectroscopy Inductively Coupled Plasma—Mass Spectrometry	200.7	22
hloride		300.0	25
		D4327-91	19
		4500-Cl ⁻ D	17
hromium	Atomic Absorption—Furnace Technique	3113 B	17
in Olliulli	Atomic Absorption—Platform	200.9	22
	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
		3120 B	17
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
olor		2120 B	17
opper	Atomic Absorption—Furnace Technique	D1688-90C	19
II.		3113 B	17
	Atomic Absorption—Platform	200.9	22
		/UU 9	,,

Contaminant	Methodology	Method	Reference
	(if appropriate)	(if appropriate)	
Copper (cont.)	Atomic Absorption—Direct Aspiration	D1688-90A	19
- of f		3111 B	17
	Inductively Coupled Plasma	200.7	22
	inductively coupled I lasma	3120 B	17
	Industrials Counted Discuss Moss Consequence		
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
Cyanide	Manual distillation followed by—	4500-CN- C	17
	1. Amenable Spectrophotometric	D2036-91B	19
	• •	4500-CN- G	17
	2. Spectrophotometric—Manual	D2036-91A	19
	2. Speed op not on the name of	4500-CN ⁻ E	17
	2. Spectrophotometrie	1-3300-85	24
	3. Spectrophotometric—	225.4	25
	Manual—Semiautomated	335.4	25
	4. Selective Electrode	4500-CN ⁻ F	17
Fluoride	Colorimetric SPADNS, with distillation	4500-F B&D	17
	Ion Chromatography	300.0	25
		D4327-91	19
		4110B	17
	Manual Electrode	D1179-93B	19
	Manual Electrode		
	Automated Alizarin Fluoride Blue, with	4500-F- C	17
	distillation (complexone)	4500-F E	17
	distination (complexone)		
	A constant of the state of the	129-71W	9
	Automated Ion Selective Electrode	380-75WE	10
Foaming Agents		5540 C	17
Iron		200.7	22
non		200.9	22
		3120 B	17
		3111 B	17
		3113 B	17
Lead	Atomic Absorption—Furnace Technique	D3559-90D	19
		3113 B	17
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
	Atomic Absorption—Platform Furnace	200.9	22
Managari		200.7	22
Manganese		200.7	22
		200.8	22
		200.9	22
		3120 B	17
		3111 B	17
		3113 B	17
Mercury	Manual cold vapor technique	245.1	22
IVICICUI y	ivianuai coru vapor tecinnique		
		D3223-91	19
		3112 B	17
	Automated cold vapor technique	245.2	1
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
Nickel	Atomic Absorption—Direct Aspiration	3111 B	17
TICKCI			
	Atomic Absorption—Platform Technique	200.9	22

Contaminant	Methodology	Method	Reference
	(if appropriate)	(if appropriate)	
lickel (cont.)	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
viekei (coiit.)	mudetivery coupled Flasina—Emission Spectroscopy	3120 B	17
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
	Atomic Absorption—Graphite Furnace	3113 B	17
litrate	Manual Cadmium Reduction	D3867-90B	19
		$4500-NO_3^-E$	17
	Automated Cadmium Reduction	353.2	25
	Automated Cadmidin Reduction	D3867-90A	19
		4500-NO ₃ -F	17
	Ion Selective Electrode	4500-NO ₃ -D	17
	ion selective Electrode	2	
	T Ol 1	601	26
	Ion Chromatography	300.0	25
		B1011	8
		4110 B	17
		D4327-91	19
litrite	Spectrophotometric	4500-NO ₃ -B	17
· -	Automated Cadmium Reduction	353.2	25
	Automated Cadmidin Reduction	D3867-90A	
			19
		$4500-NO_3^-F$	17
	Manual Cadmium Reduction	D3867-90B	19
		4500-NO ₃ -E	17
	Ion Chromatography	300.0	25
	ion emonutegraphy	B1011	8
		D4327-91	19
		4110 B	17
Odor		2150 B	17
Operational Monitoring—			17
General			17
perational Monitoring—			
pН	pH Value	150.1	1
		150.2	1
		D1293-84	19
		4500-H ⁺ B	17
Operational Monitoring—			
Residual Disinfectant			
Monitoring—	A company of the Comp	4500 010 0	17
Chlorine Dioxide	Amperometric Titration	4500 -ClO $_2$ C	17
		4500-ClO ₂ E	17
	DPD Method	4500-ClO ₂ D	17
Operational Monitoring— Residual Disinfectant		2	
Monitoring—			
Free Chlorine	Amperometric Titration	4500-Cl D	17
	DPD Ferrous Titrimetric	4500-Cl F	17
	DPD Colorimetric	4500-Cl G	17
	Syringaldazine (FACTS)	4500-Cl H	17
Combined			
G1 1 1	Amperometric Titration	4500-C1 D	17
Chlorine	Amperometric Titration—low level measurement	4500-Cl E	17
Chlorine	rumperometric ritiation for level measurement		
Chlorine	DPD Ferrous Titrimetric	4500-Cl F	17
Chlorine			17 17



Contaminant	Methodology	Method	Reference
	(if appropriate)	(if appropriate)	
Onavetional Manitarina			
Operational Monitoring— Lead and Copper			
Regulation	pH Electrometric	150.1	1
Regulation	pri Electrometre	150.2	1
		D1293-84	19
		4500-H ⁺ B	17
	Conductivity	D1125-91A	19
	Conductivity	2510 B	17
	Calcium (EDTA Titrimetric)	D511-93A	19
	Calcium (EDIA Turmetre)	3500-Ca D	17
	(Inductively Coupled Plasma)	200.7	22
	(mudchvery Coupled Flashia)	3120 B	17
	(Atomic Abcorntion Direct Achirotion)	D511-93B	19
	(Atomic Absorption—Direct Aspiration)	3111 В	
	Alleslinites (Titulos atuis)		17
	Alkalinity (Titrimetric)	D1067-92B	19
		2320 B	17
	(Electrometric Titration)	I-1030-85	24
	Orthophosphate (unfiltered, no digestion		
	or hydrolysis)		
	(Colorimetric, automated ascorbic acid)	365.1	25
		4500-P F	17
	(Colorimetric, ascorbic acid		
	single reagent)	D515-88A	19
		4500-P E	17
	(Colorimetric, phosphomolybdate,		
	automated—segmented flow,		
	automated discrete)	I-1601-85	24
		I-2601-90	24
		I-2598-85	24
	(Ion Chromatography)	300.0	25
		D4327-91	19
		4110	17
	Silica (Colorimetric, molybdate		
	blue, automated—segmented flow)	I-1700-85	24
		I-2700-85	24
	(Colorimetric)	D859-88	19
	(Molybdosilicate)	4500-Si D	17
	(Heteropoly blue)	4500-Si E	17
	(Automated method for		
	molybdate—reatine silica)	4500-Si F	17
	(Inductively Coupled Plasma)	200.7	22
		3120 B	17
		2120 2	
perational Monitoring—			
Residual Disinfectant			
Monitoring—			
Ozone	Indigo	4500-O B	17
Ozone	maigo	1300 G B	17
perational Monitoring—			
	Thermometric	2550 B	17
Temperature	THETHIOHIEUTE	2330 B	17
a di amuali da a			11
adionuclides			11
			12
	T	700	13
	Examination of Water & Wastewater for Radioactivity	700	3

Contaminant	Methodology	Method	Reference
	(if appropriate)	(if appropriate)	
adionuclides (cont.)	Gamma Spectrometry in Water	D2459	2
	Microquantities of Uranium in		
	Water by Fluorometry	D2907	2
econdary Contaminants			3
elenium	Atomic Absorption—Hydride Generation	D3859-93A	19
		3114 B	17
	Atomic Absorption—Graphite Furnace	D3859-93	19
		3113 B	17
	Atomic Absorption—Platform	200.9	22
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
ilver	Atomic Absorption—Graphite Furnace	I-3720-85	24
	Inductively Coupled Plasma—Emission Spectroscopy	200.7	22
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
	Atomic Absorption—Platform Technique	200.9	22
		3120 B	17
		3111 B	17
		3113 B	17
ulfate	Colorimetric—Methylthymol blue	375.2	25
	Gravimetric	4500-SO ₄ ²⁻ C, D	17
	Turbidimetric	4500-SO ₄ ²⁻ F	17
	Ion Chromatography	300.0	25
		4110	17
		D4327-91	19
nallium	Atomic Absorption—Platform Technique	200.9	22
	Inductively Coupled Plasma—Mass Spectrometry	200.8	22
otal Dissolved olids		2540 C	17
urbidity	Nephelometric	2130 B	17
,	1	180.1	25
	Great Lakes Instruments	Method 2	18
odium	Inductively Coupled Plasma	200.7	22
	Atomic Absorption—Direct Aspiration	3111 B	17
inc		200.8	22
		3111 B	17
		3111 B	1 /

- (A) References for analytical methods in 10 CSR 60-5.010(1):
- 1. "Methods of Chemical Analysis of Water and Wastes," EPA Environmental Monitoring and Support Laboratory, Cincinnati, OH 45268 (EPA-600/4-79-020), March 1983. Available from National Technical Information Service, PB84-128677. Methods 150.1, 150.2 and 246.2 are also available from U.S. EPA, EMSL, Cincinnati, OH 45268.
- 2. Annual Book of ASTM Standards, Vols. 11.01 and 11.02, 1991, American Society for Testing and Materials, 1916 Race Street, Philadelphia, PA 19103.
- 3. Standard Methods for the Examination of Water and Wastewater, 16th edition, American Public Health Association, American Water Works Association, Water Pollution Control Federation, 1985.
- 4. "Methods for Determination of Inorganic Substances in Water and Fluvial Sediments," *Techniques of Water-Resources Investigations of the U.S. Geological Survey Books*, Book 5, Chapter A1, Third Edition, 1989. Available at Superintendent of Documents, U.S. Government Printing Office, Washington, DC 20402.
 - 5. "Orion Guide to Water and Wastewater Analysis." Form WeWWG/5880, p. 5, 1985. Orion Research, Inc., Cambridge, MA 02139.
- 6. 200.7A "Inductively Coupled Plasma Atomic Emission Analysis of Drinking Water," Appendix to Method 200.7, March 1987, U.S. EPA, Environmental Monitoring and Support Laboratory, Cincinnati, OH 45268.

- 7. "Analytical Method for Determination of Asbestos Fibers in Water," Method 100.1, EPA-600/4-83-043, September 1983. Available from National Technical Information Service, PB83-260471.
- 8. "Waters Test Method for the Determination of Nitrite/Nitrate in Water Using Single Column Ion Chromatography," Method B-1011, Millipore Corporation, Waters Chromatography Division, 34 Maple Street, Milford, MA 01757.
- 9. "Fluoride in Water and Wastewater," Industrial Method 129-71W, Technicon Industrial Systems, Tarrytown, NY 10591, December 1972.
- 10. "Fluoride in Water and Wastewater," Industrial Method No. 380-75WE, Technicon Industrial Systems, Tarrytown, NY 10591, February 1976.
- 11. "Radiochemical Methodology for Drinking Water," Environmental Monitoring Support Laboratory, EPA-600/4-75-008, U.S. EPA, Cincinnati, OH 45268.
- 12. "Procedures for Radiochemical Analysis of Nuclear Reactor Aqueous Solutions," H.L. Krieger and S. Gold, EPA-R4-730014, U.S. EPA, Cincinnati, OH, May 1973.
 - 13. HASL Procedure Manual, edited by John H. Harley, HASL 300, ERDA Health and Safety Laboratory, New York, NY, 1973.
- 14. "Determination of and Trace Elements in Water and Wastes by Inductively Coupled Plasma-Mass Spectrometry," Method 200.8, version 4.3, August 1990, EPA, Environmental Monitoring and Systems Laboratory, Cincinnati, OH 45268. Available from ORD Publication, CERI, EPA, Cincinnati, OH 45268.
- 15. "Determination of Metals and Trace Elements by Stabilized Temperature Graphite Furnace Atomic Absorption Spectrometry," Method 200.9, version 1.1, August 1990, EPA, Environmental Monitoring and Systems Laboratory, Cincinnati, OH 45268.
- 16. "Determination of Ozone in Water by the Indigo Method; A Submitted Standard Method," *Ozone Science and Engineering*, Volume 4, pages 169–176, Pergamon Press Ltd., 1982.
- 17. Standard Methods for the Examination of Water and Wastewater, 18th edition, American Public Health Association, American Water Works Association, Water Pollution Control Federation, 1992.
 - 18. "Turbidity," GLI Method 2, November 2, 1992, Great Lakes Instruments, Inc., 8855 North 55 Street, Milwaukee, WI 53223.
- 19. Annual Book of ASTM Standards, Vols. 11.01 and 11.02, 1994, American Society for Testing and Materials, 1916 Race Street, Philadelphia, PA 19103.
- 20. "Determination of Metals and Trace Elements in Water and Wastes by Inductively Coupled Plasma-Mass Spectrometry" Method 200.8, August 1990, Revision 3.2 EPA EMSL. Available from U.S. EPA, EMSL Cincinnati, OH 45268.
- 21. "Determination of Inorganic Ions in Water by Ion Chromatography" Method 300.8, December 1989, U.S. EPA EMSL. Available from U.S. EPA, EMSL, Cincinnati, OH 45268.
- 22. "Methods for the Determination of Metals in Environmental Samples—Supplement I, EPA-600/R-94-111, May 1994." Available from National Technical Information Service (NTIS) NTIS PB 94-184942, U.S. Department of Commerce, 5285 Port Royal Road, Springfield, VA 22161. The toll free number is (800) 553-6847.
- 23. "Method 300. Determination of Inorganic Anions in Water by Ion Chromatography." Inorganic Chemistry Branch, Environmental Monitoring Systems Laboratory, August 1991.
- 24. Available from the Books and Open-File Reports Section, United States Geological Survey, Federal Center, Box 25425, Denver, CO 80225-0425.
- 25. "Methods for the Determination of Inorganic Substances in Environmental Samples," EPA-600/R-93-100, August 1993. Available from National Technical Information Service, PB94-121811.
- 26. The procedure shall be done in accordance with the Technical Bulletin 601, "Standard Method of Test for Nitrate in Drinking Water," July 1994, PN 221890-001, Analytical Technology, Inc. Available from ATI, Orion, 529 Main Street, Boston, MA 02129.
- 27. "Determination of Asbestos Structure over 10- μm in Length in Drinking Water," Method 100.2, EPA-600/R-94-134, June 1994. Available from NTIS, PB94-201902.

Contaminant

Dinoseb

(2) Organic Contaminants. Unless substitute methods are approved, the following table includes acceptable analysis procedures for organic contaminants:

includes acceptable analysis proc		Dinoseo	515.1
organic contaminants:	edules loi		515.2
organic contaminants.			555
Gtt	M-41- J	Diquat	549.1
Contaminant	Method	Endothall	548.1
2.2.7.0 ECDD (D: :)	1612	Endrin	505
2,3,7,8-TCDD (Dioxin)	1613		508
2,4-D	515.1		525.2
	515.2		508.1
	555	Ethylene dibromide (EDB)	
2,4,5-TP (Silvex)			551
515.1515.2,	555	Glyphosate	547
3-Hydrodroxycarbofuran	531.1		6651
	6610	Heptachlor	505
Alachlor	505		508
	507		525.2
	525.2		508.1
	508.1	Heptachlor epoxide	505
Aldicarb	531.1		508
	6610		525.2
Aldicarb sulfoxide	531.1		508.1
	6610	Hexachlorobenzene	505
Aldicarb sulfone	531.1		508
Thereard surrone	6610		525.2
Aldrin	505		508.1
Aldrin	508	Hexachlorocyclopentadiene	
	525.2		525.2
			508
A	508.1		508.1
Atrazine	505	Lindane	505
	507		508
	525.2		525.2
	508.1		508.1
Benzo(a)pyrene	525.2	Methomyl	531.1
	550	Wethomy	6610
	550.1	Methoxychlor	505
Butachlor	507	Wethoxyemor	525.2
	525.2		508.1
Carbaryl	531.1	Metolachlor	507
	6610	Wetolaemoi	508.1
Carbofuran	531.1		525.2
	6610	Metribuzin	507
Chlordane	505	Metriouzin	508.1
	508		525.2
	525.2	Oxamyl (vydate)	531.1
	508.1	Oxamyi (vydate)	6610
Dalapon	515.1	Dontachlorophonol	
p	552.1	Pentachlorophenol	515.1
Di(2-ethylhexyl)adipate	506		515.2
21(2 cm) meny manpane	525.2		525.2
Di(2-ethylhexyl)phthalate	506	Distance.	555
Di(2-ethymexy)/phthalate	525.2	Picloram	515.1
Dibromochloropropane (DBCP)	504.1		515.2
Dioromocinoropiopane (DBCI)	551	D 1 11 1 1 1 1 1 1	555 505
Dicamba		Polychlorinated biphenyls	505
Dicaillua	515.1 515.2		(as
	515.2		Aroclors)
Dioldrin	555 505		508
Dieldrin	505		(as
	508		Aroclors)
	508.1		508A (as deca-
	525.2		chlorobiphenyl)

Contaminant	Method
Propachlor	508
	508.1
	525.2
Simazine	505
	507
	508.1
	525.2
Toxaphene	505
-	508
	525.2
Total Trihalomethanes	502.2
	524.2
	551
Volatile Organic Chemicals	
(regulated and unregulated)	502.2
,	524.2

Footnotes

Method

515.1

- 1) A nitrogen-phosphorous detector should be substituted for the electron capture detector in Method 505 (or another approved method should be used) to determine alachlor, atrazine and simazine, if lower detection limits are required.
- 2) PCBs are qualitatively identified as Aroclors and measured for compliance purposes as decachlorobiphenyl. Each system which monitors for PCBs shall analyze each sample using either Method 505 or Method 508.
- 3) Analyses of total trihalomethanes shall be conducted in accordance with these methods and "Technical Notes on Drinking Water Methods," EPA-600/R94-173, October 1994, which is available at NTIS, PB95-104766.
 4) In addition to Methods 502.2 and 524.2, analysis for bromodichloromethane, bromoform,* chlorodibromomethane, chloroform, carbon tetrachloride, tetrachlorethylene, 1,1,1-trichloroethane, and trichloroethylene may also be conducted by EPA Method 551. Analysis for 1,2,3-trichloropropane may be conducted by Methods 502.1, 524.2 and 504.1.

References for analytical methods in 10 CSR 60-5.010(2): Methods 502.2, 505, 507, 508, 508A, 515.1 and 531.1 are in "Methods for the Determination of Organic Compounds in Drinking Water," EPA-600/4-88-039, December 1988, revised July 1991. Methods 506, 547, 550, 550.1 and 551 are in "Methods for the Determination of Organic Compounds in Drinking Water—Supplement I," EPA-600-4-90-020, July 1990. Methods 515.2, 524.2, 548.1, 549.1, 552.1 and 555 are in "Methods for the Determination of Organic Compounds in Drinking Water-Supplement II," EPA-600/R-92-129, August 1992. Method 1613 is titled "Tetra-through Octa-Chlorinated Dioxins and Furans by Isotope-Dilution HRGC/HRMS," EPA-821B-94-005, October 1994. These documents are available from National Technical Information Service (NTIS) NTIS PB91-231480, PB91-146027, PB92-207703, and PB95-104774. U.S. Department of 5285 Port Royal Road, Commerce, Springfield, VA 22161. The toll free number is (800) 553-6847. Method 6651 shall be followed in accordance with the 18th edition of "Standard Methods for the Examination of Water and Wastewater," 1992. Available from the American Public Health Association, 1015 Fifteenth Street NW, Washington, DC 20005. Method 6610 shall be followed in accordance with the "Supplement to the 18th Edition of Standard Methods for the Examination of Water and Wastewater," 1994. Available from the American Public Health Association, 1015 Fifteenth Street NW, Washington, DC 20005. EPA Methods 504.1, 508.1 and 525.2 are available from U.S. EPA EMSL, Cincinnati, OH 45268. The phone number is (513) 569-7586. Other analytical test procedures are contained in Technical Notes on Drinking Water Methods, EPA-600/R-94-173, October 1994, NTIS PB95-104766.

- 502.1 Volatile halogenated organic chemicals in water by purge and trap gas chromatography
- 502.2 Volatile organic compounds in water by purge and trap capillary column gas chromatography with photoionization and electrolytic conductivity detectors in series
- 503.1 Volatile aromatic and unsaturated organic compounds in water by purge and trap gas chromatography
- 504.1 1,2-Dibromoethane (EDB), 1,2-Dibromo-3-chloropropane (DBCP), and 1,2,3-Trichloropropane (123TCP) in Water by Microextraction and Gas Chromatography
- 505 Analysis of organohalide pesticides and commercial polychlorinated biphenyl products (Aroclors) in water by microextraction and gas chromatography
- 506 Determination of phthalate and adipate esters in drinking water by liquid-liquid extraction or liquid-solid extraction and gas chromatography with photoionization detection
- 507 Determination of nitrogen- and phosphorus-containing pesticides in groundwater by gas chromatography with a nitrogen-phosphorus detector
- 508 Determination of chlorinated pesticides in water by gas chromatography with an electron capture detector

- 508.1 Determination of chlorinated pesticides, herbicides, and organohalides by liquid-solid extraction and electron capture gas chromatography
- 508A Screening for polychlorinated biphenyls by perchlorination and gas chromatography (for quantification if detected with Method 505 or 508)
- 515.1 Determination of chlorinated acids in water by gas chromatography with an electron capture detector, revision 5.0, May 1991
- 524.1 Measurement of purgeable organic compounds in water by purged column gas chromatography/mass spectrophotometry
- 524.2 Volatile organic chemicals in water by purge and trap capillary column gas chromatography/mass spectrophotometry
- 525.2 Determination of organic compounds in drinking water by liquid-solid extraction in capillary column gas chromatography/mass spectrometry
- 531.1 Measurement of N-methyl carbamoyloximes and N-methyl carbamates in water by direct aqueous injection HPLC with post-column derivatization
- 547 Analysis of glyphosate in drinking water by direct-aqueous-injection HPLC, with post-column derivatization
- 548 Determination of endothall in aqueous samples
- 549.1 Determination of diquat and paraquat in drinking water by liquid-solid extraction and high performance liquid chromatography with ultraviolet detection
- 550 Determination of polycyclic aromatic hydrocarbons in drinking water by liquid-liquid extraction and HPLC with coupled ultraviolet and fluorescence detection
- 550.1 Determination of polycyclic aromatic hydrocarbons in drinking water by liquid-solid extraction and HPLC with coupled ultraviolet and fluorescence detection
- 552 Determination of polycyclic aromatic hydrocarbons in drinking water by liquid-solid extraction and HPLC with coupled ultraviolet and fluorescence detection
- 551 Determination of chlorination disinfection byproducts and chlorinated solvents in drinking water by liquidliquid extraction and gas chromatography with electron-capture detection

- 555 Determination of chlorinated acids in water by high performance liquid chromatography with a photodiode array ultraviolet detector
- 1613 "Tetra-through Octa-Chlorinated Dioxins and Furans by Isotope Dilution." This method is available from U.S. EPA-OST, Sample Control Center, P.O. Box 1407, Alexandria, VA 22313.
- 6610 Carbamate pesticides 6651 Glyphosate herbicide
- (3) Microbiological Contaminants. Unless substitute methods are approved, this section (3) lists acceptable analysis procedures for microbiological contaminants.

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(A) Escherichia coli. the (E. coli) Public water systems must conduct analysis of *Escherichia coli* in accordance with one (1) of following analytical methods:

- 1. EC medium supplemented with fifty (50) μ /ml of 4-methylumbelliferyl-beta-D-glucuronide (MUG) (final concentration). EC medium is described in *Standard Methods for the Examination of Water and Wastewater*, 1992, American Public Health Association, 18th edition, Method 9221E, p. 9–52, paragraph 1a. MUG may be added to EC medium before autoclaving. EC medium supplemented with fifty (50) μ /ml of MUG is commercially available. At least ten (10) ml of EC medium supplemented with MUG must be used. The inner inverted fermentation tube may be omitted. The procedure for transferring a total coliform-positive culture to EC medium supplemented with MUG shall be as specified in 10 CSR 60-5.010(1)(B)2. for transferring a total coliform-positive culture to EC medium. Observe fluorescence with an ultraviolet light (366 nm) in the dark after incubating tube at 44.5 \pm 0.2 degrees Celsius for 24 \pm 2 hours;
- 2. Nutrient agar supplemented with 100 μ /ml MUG (final concentration). Nutrient agar is described in *Standard Methods for the Examination of Water and Wastewater*, 1992, American Public Health Association, 18th edition, p. 9-47 to 9-48. This test is used to determine if a total coliform-positive sample, as determined by the membrane filter technique (MFT) or any other method in which a membrane filter is used, contains *E. coli*. Transfer the membrane filter containing a total coliform colony(ies) to nutrient agar supplemented with 100 μ /ml (final concentration) MUG. After incubating the agar plate at thirty-five degrees Celsius (35°C) for four (4) hours, observe the colony(ies) under ultraviolet light (366 nm) in the dark for fluorescence. If fluorescence is visible, *E. coli* are present;
- 3. Minimal Medium ONPG-MUG (MMO-MUG) Test, as set forth in the article "National Field Evaluation of a Defined Substrate Method for the Simultaneous Detection of Total Coliforms and *Escherichia coli* from Drinking Water: Comparison with Presence-Absence Techniques" (Edberg et al.), Applied and Environmental Microbiology, Volume 55, pp. 1003–1008, April 1989. (Note: The Autoanalysis Colilert System is an MMO-MUG test.) If the MMO-MUG test is total coliform-positive after a 24-hour incubation, test the medium for fluorescence with a 366-nm ultraviolet light (preferably with a 6-watt lamp) in the dark. If fluorescence is observed, the sample is *E. coli*-positive. If fluorescence is questionable (cannot be definitively read) after 24 hours incubation, incubate the culture for an additional four (4) hours (but not to exceed 28 hours total). And again test the medium for flourescence. The MMO-MUG Test with hepes buffer in lieu of phosphate buffer is the only approved formulation for the detection of *E. coli*;
- 4. As an option to paragraph (3)(A)3. of this rule, a system with a total coliform-positive, MUG-negative, MMO-MUG test may further analyze the culture for the presence of *E. coli* by transferring a 0.1 ml 28-hour MMO-MUG culture to EC Medium + MUG. Observation of the results are described in 40 CFR 141.21(f)(6)(i); or
- 5. The Colisure test. A description of the Colisure test may be obtained from the Millipore Corporation, Technical Services Department, 80 Ashby Road, Bedford, MA 01730.
- 1. Public water systems must conduct fecal coliform analysis in accordance with the following procedure: When the multiple-tube fermentation (MTF) technique or presence-absence (P-A) coliform test is used to test for total coliform, shake the lactose-positive presumptive tube or P-A vigorously and transfer the growth with a sterile three millimeter (3 mm) loop or sterile applicator stick into brilliant green lactose bile broth and EC medium to determine the presence of total and fecal coliforms, respectively. For EPA-approved analytical methods which use a membrane filter, transfer the total coliformpositive culture by one (1) of the following methods: Remove the membrane containing the total coliform colonies from the substrate with a sterile forceps and carefully curl and insert the membrane into a tube of EC medium (the laboratory may first remove a small portion of selected colonies for verification), swab the entire membrane filter surface with a sterile cotton swab and transfer the inoculum to EC medium (do not leave the cotton swab in the EC medium), or inoculate individual total coliformpositive colonies into EC medium. Gently shake the inoculated tubes of EC medium to insure adequate mixing and incubate in a waterbath at 44.5 \pm 0.2 degrees Celsius for 24 \pm 2 hours. Gas production of any amount in the inner fermentation tube of the EC medium indicates a positive fecal coliform test. The preparation of EC medium is described in Standard Methods for the Examination of Water and Wastewater, 1992, American Public Health Association, 18th edition, Method 9221E, p. 9-2, paragraph 1a. Public water systems need only determine the presence or absence of fecal coliforms; a determination of fecal coliform density is not required. Only this method for fecal coliform is allowed for compliance with 10 CSR 60-4.020(5); or
- 2. Standard Methods for the Examination of Water and Wastewater, 1992, American Public Health Association, 18th edition.
- A. Fecal coliform most probable number (MPN) procedures, method 9221E, pages 9-52 to 9-53 (Note: A-1 Broth may be held up to three (3) months in a tightly closed screwcap tube at four degrees Celsius (4°C):
 - B. Fecal coliform membrane filter procedure, method 9222D, pages 9-60 to 9-61.

(B) Fecal Coliform.

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(C) Heterotrophic Bacteria.

(D) Total Coliform.

Standard Methods for the Examination of Water and Wastewater, 1992, American Public Health Association, 18th edition, pour plate method, method 9215B. The time from sample collection to initiation of analysis may not exceed twenty-four (24) hours.

- 1. The standard sample volume required for total coliform analysis, regardless of analytical method used, is one hundred milliliters (100 ml). The time from sample collection to initiation of analysis may not exceed forty-eight (48) hours. If the laboratory analyzes samples after thirty (30) hours and up to forty-eight (48) hours from sample collection, the laboratory shall indicate on the report of the analysis results that the data may be invalid because of excessive delay before sample processing.
- 2. Standard Methods for the Examination of Water and Wastewater, 1992, American Public Health Association, 18th edition—
 - A. Fermentation technique, method 9221A, B.
- (I) Lactose broth, as commercially available, may be used in lieu of lauryl tryptose broth, if the system conducts at least twenty-five (25) parallel tests between this medium and lauryl tryptose broth using the water normally tested, and this comparison demonstrates that the false-positive rate for total coliforms, using lactose broth, is less than ten percent (10%).
- (II) If inverted tubes are used to detect gas production, the media should cover these tubes at least one-half (1/2) to two-thirds (2/3) after the sample is added.
- (III) No requirement exists to run the completed phase on ten percent (10%) of all total coliform-positive confirmed tubes.
 - B. Membrane filter (MF) technique, method 9222A, B, C.
 - C. Presence-absence (P-A) coliform test, method 9221D.
- (I) Six-times formulation strength may be used if the medium is filter-sterilized rather than autoclaved.
- (II) No requirement exists to run the completed phase on ten percent (10%) of all total coliform-positive confirmed tubes.
 - D. ONPG-MUG Test (also known as the Autoanalysis Colilert System, method 9223).
- E. Colisure test. The Colisure test must be incubated for twenty-eight (28) hours before examining the results. If an examination of the results at twenty-eight (28) hours is not convenient, then results may be examined at any time between twenty-eight (28) and forty-eight (48) hours.

A description of the Colisure test may be obtained from the Millipore Corporation, Technical Service Department, 80 Ashby Road, Bedford, MA 01730.

(4) Sample collection for the contaminants listed in this rule must be conducted using the sample preservation, container and maximum holding time procedures specified in the following table. All other samples for contaminants in 10 CSR 60-5.010 shall be collected in accordance with procedures contained in the appropriate analytical method.

			Holding
Contaminant	Preservative	Container	Time
Antimony	Concentrated HNO ₃ to pH < 2	P or G	6 months
Asbestos	Cool to 4°C	P or G	
Barium	Concentrated HNO ₃ to pH < 2	P or G	6 months
Beryllium	Concentrated HNO ₃ to pH < 2	P or G	6 months
Cadmium	Concentrated HNO ₃ to $pH < 2$	P or G	6 months
Chromium	Concentrated HNO ₃ to pH < 2	P or G	6 months
Copper	3 -		
Preserved	Concentrated HNO ₃ to pH < 2	P or G	6 months
Unpreserved	NONE	P or G	14 days
Cyanide	Cool to 4° C, NaOH to pH > 12	P or G	14 days
Fluoride	NONE	P or G	1 month
Lead			
Preserved	Concentrated HNO ₃ to pH < 2	P or G	6 months
Unpreserved	NONE	P or G	14 days
Mercury	Concentrated HNO ₃ to pH $<$ 2	P or G	28 days

Haldina

			Holaing
Contaminant	Preservative	Container	Time
Nickel	Concentrated HNO ₃ to pH $<$ 2	P or G	6 months
Nitrate			
Chlorinated	Cool to 4°C	P or G	28 days
Nonchlorinated	Concentrated H_2SO_4 to pH < 2	P or G	14 days
Nitrite	Cool to 4°C	P or G	48 hours
Selenium	Concentrated HNO ₃ to pH $<$ 2	P or G	6 months
Thallium	Concentrated HNO ₃ to pH < 2	P or G	6 months

- (A) If HNO₃ cannot be used because of shipping restrictions, sample may be initially preserved by icing and immediately shipping to the laboratory. Upon receipt in the laboratory, the sample must be acidified with concentrated HNO₃to pH < 2 and held for sixteen (16) hours before analysis. At time of analysis, sample container should be thoroughly rinsed with 1:1 HNO₃; washings should be added to the sample.
 - (B) P = Plastic, hard or soft.
 - (C) G = Glass, hard or soft.
- (D) In all cases samples should be analyzed as soon after collection as possible.
- (E) For cyanide see method(s) for the information for preservation.
- (F) The system shall take each sample at the same sampling point unless conditions make another sampling point more representative of each source or treatment plant.
- (G) If a system draws water from more than one (1) source and the sources are combined before distribution, the system must sample at an entry point to the distribution system during periods of normal operating conditions (that is, when water is representative of all sources being used).
- (5) The department may reduce the total number of samples a system must analyze by allowing the use of compositing. Composite samples from a maximum of five (5) sampling points are allowed provided that the detection limit of the method used for analysis is less than one-fifth (1/5) of the MCL. For a lower number of composited samples the allowable relationship between the detection limit and MCL will change proportionally. Compositing of samples must be done in the laboratory and the composite sample must be analyzed within fourteen (14) days of collection. If the population served by the system is greater than thirty-three hundred (>3,300) persons, then compositing is permitted only at sampling points within a single system. In systems serving less than or equal to thirty-three hundred ($\leq 3,300$) persons, the department may permit compositing among different systems provided the five (5)-sample limit is maintained.
- (A) Compositing of samples may be allowed for inorganic chemicals (IOCs) listed

- in 10 CSR 60-4.030(1) and synthetic organic chemicals (SOCs) listed in 10 CSR 60-4.040 and volatile organic chemicals (VOCs) listed in 10 CSR 60-4.100(2) and unregulated organic and inorganic chemicals listed in 10 CSR 60-4.110(2).
- (B) If the concentration in the composite sample is greater than or equal to 0.0005 mg/l for any organic contaminant listed in 10 CSR 60-4.100(2), then a follow-up sample must be taken within fourteen (14) days at each sampling point included in the composite. These samples must be analyzed for the contaminants which exceeded 0.0005 mg/l in the composite sample. Resampling is not required for unregulated organic and inorganic chemicals listed in 10 CSR 60-4.110(2).
- (C) If the concentration in the composite sample detects one (1) or more synthetic organic contaminants listed in subsection (6)(B) of this rule, then a follow-up sample must be taken within fourteen (14) days at each sampling point included in the composite. The follow-up sample must be analyzed for the contaminant(s) detected.
- (D) If duplicates of the original sample from each sampling point used in the composite are available, the system may use these duplicates instead of resampling. The duplicate must be analyzed and the results reported to the department within fourteen (14) days of collection.
- (E) Compositing Samples Prior to Gas Chromatograph (GC) Analysis.
- 1. Add five milliliter (5 ml) or equal larger amounts of each sample (up to five (5) samples are allowed) to a twenty-five milliliter (25 ml) glass syringe. Special precautions must be made to maintain zero headspace in the syringe.
- 2. The samples must be cooled at four degrees Centigrade (4°C) during this step to minimize volatilization losses.
- 3. Mix well and draw out a five milliliter (5 ml) aliquot for analysis.
- 4. Follow sample introduction, purging and desorption steps described in the method.
- 5. If less than five (5) samples are used for compositing, a proportionately small syringe may be used.
- (F) Compositing Samples Prior to GC/Mass Spectrophotometer (MS) Analysis.

- 1. Inject five milliliter (5 ml) or equal larger amounts of each aqueous sample (up to five (5) samples are allowed) into a twenty-five milliliter (25 ml) purging device using the sample introduction technique described in the method.
- 2. The total volume of the sample in the purging device must be twenty-five milliliters (25 ml).
- 3. Purge and desorb as described in the method.
- (G) For lead and copper monitoring, composite samples from a maximum of five (5) sampling points per composite sample are allowed with prior approval of the department. The reportable value, as required in 10 CSR 60-7.020, for each of the samples in the composite is the concentration detected multiplied by the number of samples composited. If the concentration in the composite sample is less than the detection limit, the reportable value for each of the samples in the composite is the detection limit multiplied by the number of samples composited. If the ninetieth percentile concentration, calculated in accordance with 10 CSR 60-15.010(3)(C), exceeds the lead and copper action level established in 10 CSR 60-15.010(3)(A) and (B), each of the samples from which the composite was derived must be analyzed individually and reported as required in 10 CSR 60-7.020.

(6) Detection Limits.

(A) Detection limits for inorganic chemical analytical methods are the following:

Detection Limits for Inorganic Contaminants

Contaminant	Method	Detection Limit	
		(mg/l)	
Antimony	Atomic Absorption—		
	Furnace Technique	0.003	
	Atomic Absorption—		
	Platform	0.0008	
	Inductively Coupled		
	Plasma—Mass		
	Spectrometry	0.0004	
	Atomic Absorption—		
	Gaseous Hydride	0.001	
Asbestos	Transmission Electron	0.01 million	
	Microscopy	fibers per liter	



Contaminant	Method	Detection Limit	Contaminant	Method	Detection Limit	Contaminant	Detection Limit
		(mg/l)			(mg/l)		(mg/l)
		(8 /	Nitrite	Spectrophotometric		Aroclor 1260	0.0002
Barium	Atomic Absorption—			Automated Cadmiu		Simazine	0.00007
	Furnace Technique	0.002		Reduction Manual Cadmium	0.05	Toxaphene	0.001
	Atomic Absorption— Direct Aspiration	0.1		Reduction	0.01		
	Inductively Coupled	0.1		Ion Chromatograph		(C) The detection lim	it for VOCs is 0.0005
	Plasma	0.002				mg/l.	
	Inductively Coupled		Selenium	Atomic Absorption		(D) Detection limits	for radiological con-
	Plasma (EPA method	0.001		Furnace Technique Atomic Absorption		taminants are the follow	-
	200.7)	0.001		Gaseous Hydride	0.002	turminants are the renew	
Beryllium	Atomic Absorption—			-		Radionuclide	Detection Limit
•	Furnace Technique	0.0002	Thallium	Atomic Absorption			Detection Limit
	Atomic Absorption—			Furnace Technique Atomic Absorption		Radium 226, 228	1 pCi/l
	Platform	0.00002		Platform	0.0007	Gross Alpha	3 pCi/l
	Inductively Coupled Plasma	0.0003		Inductively Couple			
	Inductively Coupled	0.0003		Plasma—Mass		Tritium	1000 pCi/l
	Plasma—Mass			Spectrometry	0.0003	Strontium-89	10 pCi/l
	Spectrometry	0.0003	(D) D	1 6 6	100 1 61	Strontium-90	2 pCi/l
Cadmium	A +		, ,	ction limits for S	OCs are the fol-	Iodine-131	1 pCi/l
Cadmium	Atomic Absorption— Furnace Technique	0.0001	lowing:			Cessium-134	10 pCi/l
	Inductively Coupled	0.0001	a		T T	Gross Beta	4 pCi/l
	Plasma	0.001	Contamina	nt	Detection Limit	Other radionuclides	1/10 of the
					(mg/l)		acceptable
Chromium	Atomic Absorption—	0.001		OD (Dioxin)	0.000000005		limit
	Furnace Technique Inductively Coupled	0.001	2,4-D		0.0001		
	Plasma	0.007	2,4,5-TP (S	ilvex)	0.0002	AUTHORITY: section	ns 640.100 and
	Inductively Coupled		Alachlor		0.0002	640.125.1, RSMo Supp.	
	Plasma (EPA Method		Aldicarb		0.0005	filed May 4, 1979, effe	
	200.7A)	0.001	Aldicarb sul	lfoxide	0.0005	Amended: Filed April 14	•
Copper	All Methods Except		Aldicarb sul	lfone	0.0008	•	00
Copper	Atomic Absorption	0.0002	Atrazine		0.0001	11, 1981. Amended: I	
	Atomic Absorption		Benzo(a)pyr	ene	0.00002	effective Aug. 31, 1988.	
	With Direct Aspiration	0.0002	Carbofuran		0.0009	opted: Filed Dec. 4, 19	
a	This was a second of the secon		Chlordane		0.0002	1991. Rescinded and red	1
Cyanide	Distillation, Spectrophoto- metric (screen)	0.02	Dalapon		0.001	31, 1992, effective Dec	3, 1992. Amended:
	Distillation, Automated	0.02	Dibromochl	oronronane	0.001	Filed Aug. 4, 1992, eff	fective May 6, 1993.
	Spectrophoto-		(DBCP)	огоргорине	0.00002	Amended: Filed May 4,	1993, effective Jan.
	metric (screen)	0.005	Di(2-ethylhe	evyl)adinate	0.0006	13, 1994. Amended: 1	Filed Feb. 1, 1996,
	Distillation, Selective	0.02		exyl)phthalate	0.0006	effective Oct. 30, 1996.	Amended: Filed July
	Electrode (screen) Distillation, Amenable,	0.02	Di(2-etilying	exyr)phinalate	0.0003	1, 1999, effective March	i 30, 2000.
	Spectrophotometric						
	(free)	0.02	Diquat		0.0004	*Original authority: 640.100 R	SMo, 1939, amended 1978,
			Endothall		0.009	1981, 1982, 1988, 1989, 1992	, 1993, 1995, 1996, 1998,
Lead	All Methods	0.001	Endrin		0.00001	1999; and 640.125, RSMo 197	78, amended 1988.
Mercury	Manual Cold Vapor			promide (EDB)	0.00001		
Mercury	Technique	0.0002	Glyphosate		0.006		
	Automated Cold Vapor		Heptachlor		0.00004	10 CSR 60-5.020 Labo	ratory Certification
	Technique	0.0002	Heptachlor		0.00002		
Nicologi	A + A 1		Hexachlorob	penzene	0.0001	PURPOSE: This rul	e establishes that
Nickel	Atomic Absorption— Furnace Technique	0.001	Hexachloroc	cyclopentadiene	0.0001	required analyses must	be done by laborato-
	Atomic Absorption—	0.001	Lindane		0.00002	ries certified by the dep	artment.
	Platform	0.0006	Methoxychle	or	0.0001		
	Inductively Coupled		Oxamyl (Vy		0.002	(1) For the purpose of	determining compli-
	Plasma	0.005	Pentachloro	phenol	0.00004	ance with this chapter,	analytical results will
	Inductively Coupled Plasma—Mass		Picloram	•	0.0001	be acceptable only if th	
	Spectrometry	0.0005		ated biphenyls (F		analyzed by a laborate	
	• •		-	lorobiphenyl)	0.0001	department.	. ,
Nitrate	Manual Cadmium	6.64	Aroclor 1		0.0001	asparament.	
	Reduction	0.01	Aroclor 1		0.00008	(2) To receive approval	to conduct analyses
	Automated Hydrazine Reduction	0.01	Aroclor 1		0.005	for antimony, asbestos,	
	Automated Cadmium	0.01	Aroclor 1		0.0003	cadmium, chromium, cy	
	Reduction	0.05			0.0003		
	Ion Selective Electrode	1	Aroclor 1			cury, nickel, nitrate, n	
	Ion Chromatography	0.01	Aroclor 1	.434	0.0001	thallium the laboratory	must—

- (A) Analyze performance evaluation samples which include those substances provided by the Environmental Protection Agency (EPA) Environmental Monitoring and Support Laboratory or equivalent samples provided by the department; and
- (B) Achieve quantitative results on the analyses that are within the following acceptance limits:

Acceptance Limit
$\pm 30\%$ at ≥ 0.006 mg/l
2 standard deviations
based on study statistics
$\pm 15\%$ at ≥ 0.15 mg/l
$\pm 15\%$ at ≥ 0.001 mg/l
$\pm 20\%$ at ≥ 0.002 mg/l
$\pm 15\%$ at ≥ 0.01 mg/l
$\pm 10\%$ at ≥ 0.50 mg/l
$\pm 10\%$ at \geq to 10 mg/l
$\pm 30\%$ at ≥ 0.005 mg/l
$\pm 30\%$ at ≥ 0.0005 mg/l
$\pm 15\%$ at ≥ 0.01 mg/l
$\pm 10\%$ at ≥ 0.4 mg/l
$\pm 15\%$ at ≥ 0.4 mg/l
$\pm 20\%$ at ≥ 0.01 mg/l
$\pm 30\%$ at ≥ 0.002 mg/l

- (3) To receive certification to conduct analyses for the contaminants in 10 CSR 60-4.100(2)(A)2-7. and (B)1.-13., the laboratory must— $\frac{1}{2}$
- (A) Analyze performance evaluation samples which include these substances provided by EPA Environmental Monitoring and Support Laboratory or equivalent samples provided by the department:
- (B) Achieve the quantitative acceptance limits in subsections (3)(C) and (D) of this rule for at least eighty percent (80%) of the regulated organic chemicals listed in 10 CSR 60-4.100(2)(A)2.-7. and (B)1.-13.;
- (C) Achieve the quantitative results on the analyses performed under subsection (3)(A) of this rule that are within plus or minus twenty percent ($\pm 20\%$) of the actual amount of the substances in the performance evaluation sample when the actual amount is greater than or equal to 0.010 mg/l;
- (D) Achieve quantitative results on the analyses performed under subsection (3)(A) of this rule that are within plus or minus forty percent ($\pm 40\%$) of the actual amount of the substances in the performance evaluation sample when the actual amount is less than 0.010 mg/l; and
- (E) Achieve a method detection limit of $0.0005 \ mg/l$.

- (4) To receive certification for vinyl chloride, the laboratory must—
- (A) Analyze performance evaluation samples provided by EPA Environmental Monitoring and Support Laboratory or equivalent samples provided by the department;
- (B) Achieve quantitative results on the analyses performed under subsection (4)(A) of this rule that are within plus or minus forty percent ($\pm 40\%$) of the actual amount of vinyl chloride in the performance evaluation sample;
- (C) Achieve a method detection limit of 0.0005 mg/l; and
- (D) Obtain certification for the contaminants listed in 10 CSR 60-4.100(2)(A)2.-7. and (B)1.-13.
- (5) To receive certification to conduct analyses for the contaminants in 10 CSR 60-4.040(1), the laboratory must—
- (A) Analyze performance evaluation samples which include those substances provided by EPA Environmental Monitoring and Support Laboratory or equivalent samples provided by the department.
- (B) Achieve quantitative results on the analyses that are within the following acceptance limits:

Contaminant	Acceptance Limit
	(percent)
2,3,7,8-TCDD (Dioxin)	2 standard
2,4-D	± 50
2,4,5-TP	± 50
Alachlor	± 45
Aldicarb	2 standard
	deviations
Aldicarb sulfoxide	2 standard
	deviations
Aldicarb sulfone	2 standard
	deviations
Atrazine	± 45
Benzo(a)pyrene	2 standard
	deviations
Carbofuran	± 45
Chlordane	± 45
Dalapon	2 standard
-	deviations
Dibromochloropropane	± 40
Di(2-ethylhexyl)adipate	2 standard
	deviations
Di(2-ethylhexyl)phthalate	2 standard
	deviations
Dinoseb	2 standard
	deviations
Diquat	2 standard
•	deviations
Endothall	2 standard
	deviations
Endrin	± 45

Ethylene dibromide	± 40
Glyphosate	2 standard
	deviations
Heptachlor	± 45
Heptachlor epoxide	± 45
Hexachlorobenzene	2 standard
	deviations
Hexachlorocyclopentadiene	2 standard
	deviations
Lindane	± 45
Methoxychlor	± 45
Oxamyl	2 standard
	deviations
Polychlorinated biphenyls	
(PCBs) (as decachloro-	
biphenyl)	0-200
Picloram	2 standard
	deviations
Simazine	2 standard
	deviations
Toxaphene	± 45
Pentachlorophenol	± 50

- (6) To receive approval to conduct analyses for copper and lead, the laboratory must—
- (A) Analyze performance evaluation samples which include those substances provided by EPA Environmental Monitoring and Support Laboratory or equivalent samples provided by the department;
- (B) Achieve quantitative acceptance limits for copper plus or minus ten percent $(\pm 10\%)$ of the actual amount in the performance evaluation sample when the actual amount is greater than or equal to 0.050 mg/l; lead plus or minus thirty percent $(\pm 30\%)$ of the actual amount in the performance evaluation sample when the actual amount is greater than or equal to 0.005 mg/l; and
- (C) Achieve a method detection limit of $0.001 \ \text{mg/l}.$
- (7) The department has the authority to allow the use of previously collected monitoring data for purposes of monitoring, if the data were collected and analyzed in accordance with the requirements of this rule.
- (8) All lead levels measured between the Practical Quantification Level (PQL) and Method Detection Limit (MDL) must be either reported as measured or they can be reported as one-half (1/2) the PQL (0.0025 mg/l). All levels below the lead MDL must be reported as zero (0).
- (9) All copper levels measured between the PQL and the MDL must be either reported as measured or they can be reported as one-half (1/2) the PQL (0.015 mg/l). All levels below the copper MDL must be reported as zero (0).

- (10) Operational monitoring measurements required by 10 CSR 60-4.080(3) shall be performed on-site by persons acceptable to the department.
- (11) The department will consider acceptance of analytical results from out-of-state laboratories upon written request.

AUTHORITY: section 640.100, RSMo (1994).* Original rule filed May 4, 1979, effective Sept. 14, 1979. Rescinded and readopted: Filed March 31, 1992, effective Dec. 3, 1992. Amended: Filed Aug. 4, 1992, effective May 6, 1993. Amended: Filed May 4, 1993, effective Jan. 13, 1994. Amended: Filed Feb. 1, 1996, effective Oct. 30, 1996.

*Original authority 1939, amended 1978, 1981, 1982, 1988, 1989, 1992, 1993, 1995.